# **WEST Search History**



DATE: Thursday, August 23, 2007

Set	Ouerv	Hit
<u>Name</u>		<u>Count</u>
DB=PGPB; $PLUR=YES$ ; $OP=ADJ$		
L10	19 and tefluthrin.CLM.	1
L9	18 and (hydride or borohydride or hydrogen.CLM.)	4
L8	17 and (reduce or reducing or reduced or reduction.CLM.)	4
L7	tetrafluoroterephthalate or tetrahaloterephthalate.CLM.	4
L6	tetrafluoroterephthalyl alcohol or tetrahaloterephthalyl alcohol or tetrafluorodimethylolbenzene or tetrahalodimethylolbenzene or tetrahalodialkylolbenzene.CLM.	1.
DB=PGPB, USPT, USOC, EPAB, JPAB, DWPI; PLUR=YES; OP=ADJ		
L5	14 and (pyrethoid or tefluthrin)	3
L4	11 and 13	3
L3	12 and (reduced or reducing or reduction)	18
L2	tetrafluoroterephthalate	21
L1	tetrafluoroterephthalyl alcohol or tetrafluorodimethylolbenzene	5
	Name DB=P L10 L9 L8 L7 L6 DB=P L5 L4 L3 L2	Name  DB=PGPB; PLUR=YES; OP=ADJ  L10 19 and tefluthrin.CLM.  L9 18 and (hydride or borohydride or hydrogen.CLM.)  L8 17 and (reduce or reducing or reduced or reduction.CLM.)  L7 tetrafluoroterephthalate or tetrahaloterephthalate.CLM.  tetrafluoroterephthalyl alcohol or tetrahaloterephthalyl alcohol or  L6 tetrafluorodimethylolbenzene or tetrahalodimethylolbenzene or tetrahalodialkylolbenzene.CLM.  DB=PGPB, USPT, USOC, EPAB, JPAB, DWPI; PLUR=YES; OP=ADJ  L5 14 and (pyrethoid or tefluthrin)  L4 11 and 13  L3 12 and (reduced or reducing or reduction)  L2 tetrafluoroterephthalate

END OF SEARCH HISTORY

#### => d his

(FILE 'HOME' ENTERED AT 12:14:10 ON 23 AUG 2007)

FILE 'REGISTRY' ENTERED AT 12:14:39 ON 23 AUG 2007

L1 STRUCTURE UPLOADED

L2 1 S L1

L3 33 S L1 FULL

L4 1 S L1 CSS FULL

FILE 'HCAPLUS, CHEMCATS' ENTERED AT 12:16:37 ON 23 AUG 2007

L5 1 S L4

FILE 'CASREACT' ENTERED AT 12:17:25 ON 23 AUG 2007

L6 STRUCTURE UPLOADED

L7 0 S L6

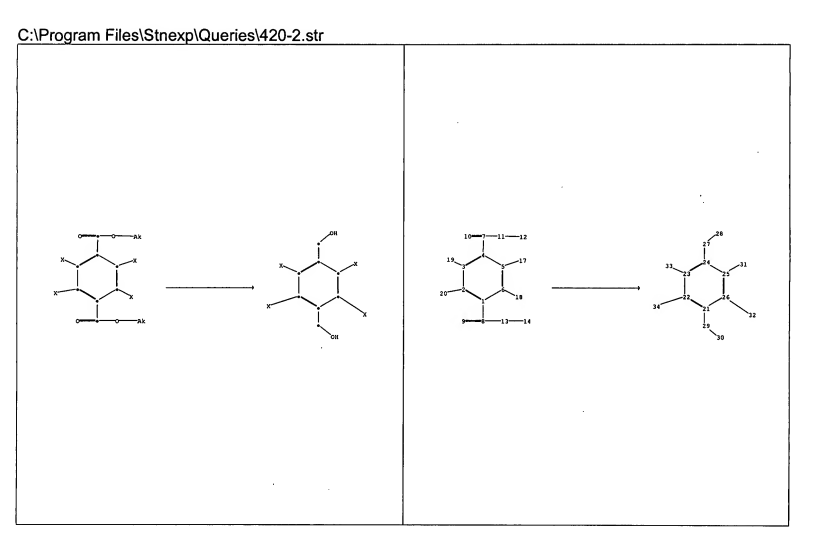
L8 3 S L6 FULL

FILE 'HCAPLUS, HCAOLD, USPATFULL, EPFULL' ENTERED AT 12:18:40 ON 23 AUG

2007

FILE 'HCAPLUS' ENTERED AT 12:19:00 ON 23 AUG 2007

L9 3 S L8



chain nodes:

7 8 9 10 11 12 13 14 17 18 19 20 27 28 29 30 31 32 33 34

ring nodes:

1 2 3 4 5 6 21 22 23 24 25 26

chain bonds:

1-8 2-20 3-19 4-7 5-17 6-18 7-10 7-11 8-9 8-13 11-12 13-14 21-29 22-34 23-33 24-27 25-31 26-32 27-28 29-30

ring bonds:

1-2 1-6 2-3 3-4 4-5 5-6 21-22 21-26 22-23 23-24 24-25 25-26

exact/norm bonds:

7-10 7-11 8-9 8-13 11-12 13-14 27-28 29-30

exact bonds:

1-8 2-20 3-19 4-7 5-17 6-18 21-29 22-34 23-33 24-27 25-31 26-32

normalized bonds:

1-2 1-6 2-3 3-4 4-5 5-6 21-22 21-26 22-23 23-24 24-25 25-26

isolated ring systems:

containing 1: 21:

## Match level:

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS8:CLASS9:CLASS10:CLASS11:CLASS 12:CLASS13:CLASS14:CLASS17:CLASS18:CLASS19:CLASS20:CLASS21:Atom 22:Atom 23:Atom 24:Atom 25:Atom 26:Atom 27:CLASS28:CLASS29:CLASS30:CLASS31:CLASS32:CLASS33:CLASS 34:CLASS

fragments assigned product role:

containing 21
fragments assigned reactant/reagent role:
 containing 1
Element Count :
 Node 12: Limited
 C,C1-7

Node 14: Limited C,C1-7

#### L1 STRUCTURE UPLOADED

=> d

L1 HAS NO ANSWERS

L1 STR

Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 12:15:01 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 4320 TO ITERATE

46.3% PROCESSED 2000 ITERATIONS

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1 ANSWERS

INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

BATCH \*\*COMPLETE\*\*

PROJECTED ITERATIONS: 82459 TO 90341

PROJECTED ANSWERS: 1 TO 131

L2 1 SEA SSS SAM L1

=> d scan

L2 1 ANSWERS REGISTRY COPYRIGHT 2007 ACS on STN

IN 1,4-Benzenedicarboxylic acid, 2,3,5,6-tetrachloro-, bimol. monoanhydride

(9CI)

MF C16 H2 C18 O7

CI COM

#### \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

#### ALL ANSWERS HAVE BEEN SCANNED

=> s l1 full FULL SEARCH INITIATED 12:15:22 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 86542 TO ITERATE

100.0% PROCESSED 86542 ITERATIONS

33 ANSWERS

SEARCH TIME: 00.00.02

L3 33 SEA SSS FUL L1

=> d scan

L3 33 ANSWERS REGISTRY COPYRIGHT 2007 ACS on STN

IN Benzoic acid, 2,3,5,6-tetrachloro-4-[[(methoxymethyl)amino]carbonyl]-,
 methyl ester (9CI)

MF C11 H9 Cl4 N O4

$$\begin{array}{c|c} \text{Cl} & \overset{\text{O}}{\parallel} \\ \text{C-NH-CH}_2\text{-OMe} \\ \\ \text{MeO-C} & \text{Cl} \\ \\ \end{array}$$

### \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):2

L3 33 ANSWERS REGISTRY COPYRIGHT 2007 ACS on STN

IN 1,4-Benzenedicarboxylic acid, 2,3,5,6-tetrachloro-, monopropyl ester (9CI)

MF C11 H8 C14 O4

$$C1$$
 $C1$ 
 $C$ 
 $C$ 
 $C$ 
 $C$ 
 $C$ 
 $C$ 

## \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

L3 33 ANSWERS REGISTRY COPYRIGHT 2007 ACS on STN

IN 1,4-Benzenedicarboxylic acid, 2,3,5,6-tetrachloro-, monoethyl ester (9CI)

MF C10 H6 Cl4 O4

CI COM

#### \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> s l1 css full\
COMBINATION OF STRUCTURE AND TEXT TERMS NOT VALID
The query entered contains both search terms created by
structure-building or screen commands and text search terms. L#s
created via the STRUCTURE or SCREEN commands must be searched in the
structures files separately from text terms or profiles. The L#
answer sets from structure searches can be used in crossover searches
and can be combined with text terms.

=> s l1 css full FULL SEARCH INITIATED 12:16:11 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 86542 TO ITERATE

100.0% PROCESSED 86542 ITERATIONS

1 ANSWERS

SEARCH TIME: 00.00.02

L4 1 SEA CSS FUL L1

=> d scan

L4 1 ANSWERS REGISTRY COPYRIGHT 2007 ACS on STN
IN Benzoic acid, 2,3,5,6-tetrafluoro-4-(hydroxymethyl)-, methyl ester (9CI)
MF C9 H6 F4 O3

$$\begin{array}{c|c} F & O \\ \hline C - OMe \\ \hline HO-CH_2 & F \end{array}$$

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

ALL ANSWERS HAVE BEEN SCANNED